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Description

This invention relates to indole derivatives, to processes for their preparation, to pharmaceutical compositions containing them and to their medical use, in particular to compounds and compositions of use in the treatment of migraine.

It has been suggested that the pain of migraine may be associated with excessive dilatation of the cranial vasculature and known treatments for migraine include the administration of compounds having vasconstrictor properties such as ergotamine. However, ergotamine is a non-selective vascoonstrictor which constricts blood vessels throughout the body and has undesirable and potentially dangerous side effects. In Migraine may also be treated by administering an analgesic usually in combination with an antiemetic but such reatments are of limited value.

More recently, indole derivatives which are selective 5HT--like receptor agonists and which exhibit selective vasconstrictor activity have been described in the art as useful in the treatment of migraine (see for example A. Doenicke, J. Brand, V. L. Perrin, Lancet, 1988, 1309-1311).

We have now found a novel group of indole derivatives which not only exhibit 5HT₁-like receptor agonist activity and selective vasoonstriction but also unexpectedly have an enhanced overall bioavailability index following administration, in particular following non-perenteral administration.

Thus the invention provides in a first aspect an indole of formula (I).

$$\mathsf{R_1R_2N-SO_2(\bar{\mathsf{CH}}_2)_2} \underbrace{}_{\substack{\mathsf{N}\\\mathsf{R_3}}}^{\mathsf{N}-\mathsf{R}}$$

30 wherein

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R₁ represents a hydrogen atom or a C₁₋₆ alkyl group;

R₂ represents a hydrogen atom or a C₁₋₆ alkyl group;

R₃ represents a hydrogen atom.

R₄ represents a hydrogen atom or a C1-3 alkyl group

35 and pharmaceutically acceptable salts and solvates (for example hydrates) thereof.

All optical isomers of compounds of general formula (I) and their mixtures including the racemic mixtures thereof are embraced by the invention.

As used herein, an alkyl group may be a straight chain (such as a methyl or ethyl) or branched chain alkyl group.

Suitable pharmacoutically acceptable salts of the indoles of general formula (i) include acid addition salts formed with organic or inorganic acids, for example, hydrochlorides, hydrobromides, sulphates, furnarates and maleates. Other salts may be useful in the preparation of compounds of formula (i), e.g. creatinine sulphates adducts.

A preferred class of compounds represented by the general formula (I) is that wherein R_1 represents a hydrogen atom or a C_{1-3} alkyl group such as a methyl group.

Another preferred class of compounds is that wherein R_2 represents a hydrogen atom or a C_{1-3} alkyl group such as methyl.

Conveniently, R₁ and R₂ together comprise from 1 to 3 carbon atoms.

The substituent R₄ is conveniently a C₁₋₃ alkyl group such as methyl.

Preferred compounds according to the invention include :-

N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide;

N.N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide:

N-Ethyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide;

N-Methyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide;

55 3-(1-Methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide;

and pharmaceutically acceptable salts and solvates thereof.

The selective 5HT₁-like receptor agonist activity and selective vasoconstrictor activity of the compounds of the invention have been demonstrated in vitro. In addition, compounds of the invention selectively

constricted the carotid arterial bed of the anaesthetised dog whilst having negligible effect on blood

Following non-parenteral, including intra-duodenal administration, the compounds of the invention show an enhanced bioavailability index in animals.

Compounds of the invention are useful in treating conditions associated with cephalic pain. In particular the compounds are useful in the treatment of migraine, cluster headache, chronic paroxysmal hemicrania and headache associated with vascular disorders and in alleviating the symptoms associated therewith.

Accordingly, the invention also provides a pharmaceutical composition which comprises at least one compound of formula (f) or a pharmaceutically acceptable salt or solvate (e.g. hydrate) thereof and formulated for administration by any convenient route. Such compositions are preferably in a form adapted for use in medicine, in particular human medicine, and can conveniently be formulated in conventional manner using one or more pharmaceutically acceptable carriers or excipients.

In a further aspect there is provided a compound of formula (I) or a salt or solvate thereof for use in therapy, in particular in human medicine. It will be appreciated that use in therapy embraces but is not a necessarily limited to use of a compound of formula (I) or a salt or solvate thereof as an active therapeutic substance.

There is also provided as a further aspect of the invention the use of a compound of formula (f) in the preparation of a medicament for use in the treatment of conditions associated with cephalic pain in particular migraine, cluster headache, chronic paroxysmal hemicrania and headache associated with a vascular disorders.

It will be appreciated that reference to treatment is intended to include prophylaxis as well as the allowards of established symptoms. Compounds according to the invention may be administered as the raw chemical but the active ingredient is preferably presented as a pharmaceutical formulation.

The active ingredient may conveniently be presented in unit dose form. A convenient unit dose formulation contains the active ingredient compound in an amount of from 0.1mg to 100mg.

The compounds according to the invention may for example be formulated for oral, sub-lingual buccal, parenteral, rectal or intranasal administration or in a form suitable for administration by inhalation or insufflation (either through the mouth or nose).

For oral administration, the pharmaceutical compositions may take the form of, for example, tablets or capsules prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g. pregelatinised maize starch, polyvinylpyrrolidone or hydroxypropyl methylcellulose); fillers (e.g. lactose, microcrystalline cellulose or calcium phosphate); lubricants (e.g. magnesium stearate, talco or silical), disintegrants (e.g. potato starch or sodium starch glycollate); or wetting agents (e.g. sodium lauryl sulphiate). The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents (e.g. sorbitol syrup, methyl cellulose or hydrogenated edible fats); emulsifying agents (e.g. lecithin or accial); non-aqueous vehicles (e.g. almond oil, oily esters or ethyl alcohol); and preservatives (e.g. methyl or propyl-phydroxybenzotates or sorbic acidi).

For buccal administration the compositions may take the form of tablets or lozenges formulated in conventional manner.

The compounds of the invention may be formulated for parenteral administration by injection, conveniently intravenous, intramuscular or subcutaneous injection, for example by bolus injection or continuous intravenous infusion. Formulations for injection may be presented in unit dosage form e.g. in ampoules or in multi-dose containers, with an added preservative.

The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilising and/or dispersing agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g. sterile ovyogen-free water, before use.

The compounds of the invention may also be formulated in rectal compositions such as suppositories or retention enemas, e.g. containing conventional suppository bases such as cocoa butter or other glyceride.

Tablets for sub-lingual administration may be formulated in a similar manner.

For intranasal administration the compounds of the invention may be used, for example, as a liquid spray, as a powder or in the form of drops.

For administration by inhalation the compounds according to the invention are conveniently delivered in the form of an aerosol spray presentation from pressurised packs or a nebuliser, with the use of a suitable propellant, e.g. dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurised aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of e.g. gelatin for use in an inhafer or insufflator may be formulated containing a powder mix of a compound of the invention and a suitable powder base 5 such as lactose or starch.

It will be appreciated that the precise dose administered will depend on the age and condition of the patient, the particular compound used and the frequency and route of administration. The compound may be administered in single or divided doses and may be administered one or more times, for example 1 to 4 times per day.

A proposed dose of the compounds of the invention for oral, sub-lingual parenteral, buccal, rectal or intransast administration to man (of approximately 70kg bodyweight) for the treatment of migraine is 0.1 to 100mg of the active ingredient per unit dose which could be administered, for example, 1 to 4 times per day.

For oral administration a unit dose will preferably contain from 2 to 50 mg of the active ingredient. A unit dose for parenteral administration will preferably contain 0.2 to 5 mg of the active ingredient.

Aerosof formulations are preferably arranged so that each metered dose or 'puff' delivered from a pressurised aerosol contains 0.2 mg to 2 mg of a compound of the invention, and capsules and catridiges delivered from an insufficient or an inhaler, contain 0.2 mg to 20 mg of a compound of the invention. The overall daily dose by inhalation with an aerosol will be within the range 1 mg to 100 mg. Administration may be several times daily, for example, from 2 to 8 times, giving for example 1, 2 or 3 doses each time.

Dosages of the compounds of the invention for rectal, sub-lingual or intranasal administration are similar to those for oral administration.

The compounds of the invention may, if desired, be administered in combination with one or more other terrapeutic agents, such as analgesics, anti-inflammatory agents and anti-nauseants, and formulated for administration by any convenient route in conventional manner. Appropriate doses will be readily appreciated by those skilled in the art.

Compounds of formula (I) and pharmaceutically acceptable salts and solvates (e.g. hydrates) thereof, may be prepared by methods known in the art for the preparation of analogous compounds. In particular the compounds of formula (I) maybe prepared by the methods outlined below and which form a further 30 aspect of the invention. In the following processes, R₁, R₂, R₃ and R₄, are as defined for formula (I) unless otherwise socified.

According to one general process (A) compounds of formula (I) may be prepared by reduction of the corresponding compounds of formula (II).

$$R_1R_2N-SO_2(CH_2)_2$$

The compounds of formula (II) are themselves novel compounds and a further part of the invention. The compounds of formula (II) have also been found to be potent and selective vasoconstrictors.

The reduction process may conveniently be carried out in the presence of hydrogen and a noble metal ocatalyst, such as palladium, Raney nickel, platinum, platinum oxide or rhodium which may be supported, for example, on charcoal. Alternatively a homogenous catalyst such as tris(triphenylphosphine) rhodium chloride may be used. The reduction may be carried out in a solvent such as an alcohol e.g. methanol or ethanol, an ether e.g. dioxan, an ester e.g. ethyl acetate or an amide e.g. dimethylformamide and conveniently at a temporarture of from -10 to +50°C.

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The compounds of formula (II) may be prepared by condensing a compound of formula (III):

or a protected or activated derivative thereof, with a piperidone of formula (IV):

or a salt or protected derivative thereof.

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The condensation reaction may be effected in a suitable reaction medium in the presence of an acid or a base, conveniently at a temperature of 25 to 120 °C.

Acids which may be employed in the above process include organic and inorganic acids such as sulphonic acids (e.g. p-toluenesulphonic acid), carboxylic acids (e.g. acetic acid) and preferably strong inorganic acids such as polyphosphoric acid, sulphuric acid and hydrochloric acid. Sulbable solvents for the reaction include inert solvents such as ethers (e.g. tetrahydrofuran or dioxan), alcohols (e.g. ethanol) and chlorinated hydrocarbons (e.g. chloroform or carbon tetrachloride). In some cases the acid may also act as the reaction solvent.

Bases which may be employed in the above process include alkali metal hydroxides (e.g. potassium hydroxide), alkali metal alkoxides (e.g. sodium or potassium methoxide, ethoxide or t- butoxide), alkali metal hydrides (e.g. sodium hydride) and alkali metal amides (e.g. sodamide). Suitable solvents for the reaction include alcohols (e.g. methanol or ethanol), ethers (e.g. tetrahydrofuran or dioxan) and dimethylsulphoxide.

Intermediates of formula (III) may be prepared by conventional methods for example by reacting an se amine of formula R-R₂NH with the 3-unsubstituted analogues of compounds of formula (V) (as described hereinafter) using the methods described for process (B) hereinafter.

According to another general process (B), a compound of formula (I) may also be prepared by condensing an amine of formula R₁ R₂ NH with an acid of formula (V)

$$HO-SO_2-(CH_2)_2$$

$$R_3$$

$$(Y)$$

or an acylating agent corresponding thereto, or a salt (for example, an organic or inorganic acid addition salt such as the hydrochloride, hydrobromide, maleate, sulphate or creatinine sulphate adduct) or a protected derivative thereof.

Acylating agents corresponding to the acid of general formula (V) which may conveniently be used in the above process include acid halides, for example sulphonyl chlorides.

The condensation process involving the acylating agents may be effected in a suitable reaction medium and conveniently at a temperature of from -70 to +150 °C. Thus the condensation reaction using an acid

halide may be effected in a suitable reaction medium such as an amide (e.g. N,N'-dimethylformamide), an ether (e.g. tetrahydrofuran), a nitrile (e.g. acetonitrile), a haloalkane (e.g. dichloromethane) or mixtures thereof, optionally in the presence of a base such as pyridine or triethylamine or an inorganic base as calcium carbonate or sodium bicarbonate.

Where it is desired to prepare a compound of formula (I) in which R1 and R2 are both hydrogen atoms, ammonia may be used in the form of aqueous ammonia or in a solvent such as methanol.

Compounds of formula (V) and acylating agents corresponding thereto are novel and as such constitute a further feature of the invention. Compounds of formula (V) or acylating agents corresponding thereto may be prepared by methods analogous to those described in UK Patent Specification 2150932 and 'A 10 Chemistry of Heterocyclic compounds - Indoles Part III, Chapter VI, edited by W. J. Houlihan (1972) Wiley Interscience, New York or by processes, such as process (A), as described herein.

According to another general process (C), a compound of formula (I) may be prepared by cyclisation of a compound of formula (VI)

$$R_1R_2NSO_2(CH_2)_2$$
 $NR_3N=CHCH_2$
 NR_4
 NR_4

The process is desirably carried out in the presence of polyphosphate ester in a reaction medium 25 which may comprise one or more organic solvents, preferably halogenated hydrocarbons such as chloroform, dichloromethane, dichloroethane, dichlorodifluoromethane, or mixtures thereof. Polyphosphate ester is a mixture of esters which may be prepared from phosphorus pentoxide, diethylether and chloroform according to the method described in 'Reagents for Organic Synthesis', (Fieser and Fieser, John Wiley and Sons 1967).

Alternatively the cyclisation may be carried out in aqueous or non-aqueous media, in the presence of an acid catalyst. When an aqueous medium is employed this may be an aqueous organic solvent such as an aqueous alcohol (e.g. methanol, ethanol or isopropanol) or an aqueous ether (e.g. dioxan or tetrahydrofuran) as well as mixtures of such solvents and the acid catalyst may be for example an inorganic acid such as concentrated hydrochloric, sulphuric or polyphosphoric acid. (In some cases the acid catalyst 35 may also act as the reaction solvent). In an anhydrous reaction medium, which may comprise one or more alcohols or ethers (e.g. as described above) or esters (e.g. ethyl acetate), the acid catalyst will generally be a Lewis acid such as boron trifluoride, zinc chloride or magnesium chloride. The cyclisation reaction may conveniently be carried out at temperatures of from 20 to 200 °C preferably 50 to 125 °C.

According to a particular embodiment of this process, compounds of formula (I) may be prepared 40 directly of the reaction of a compound of formula (VII):

$$R_1R_2NSO_2(CH_2)_2$$
 (VII)

50 or a salt thereof, with a compound of formula (VIII)

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or a salt or protected derivative thereof (such as an acetal formed, for example, with an appropriate alkylorthoformate) using the appropriate conditions as described above. It will be appreciated that in this embodiment, a compound of formula (VI) is formed as an intermediate, and may be reacted in situ to form the desired compound of ceneral formula (I).

5 Compounds of general formula (VI) may, if desired, be isolated as intermediates during the process for the preparation of compounds of formula (I) wherein a compound of formula (VII), or a salt or protected derivative thereof, is reacted with a compound of formula (VIII), or a salt or protected derivative thereof, in water or in a suitable solvent, such as an aqueous alcohol (e.g. methanol) at a temperature of, for example, 20 to 100 °C. If an acetal or ketal of a compound of formula (VIII) is used, it may be necessary to carry out 10 the reaction in the presence of an acid (for example, acid or hydrochloric acid).

Compounds of general formula (VII) may be prepared in a number of conventional steps, from compounds of formula (IX):

$$R_1R_2NSO_2(CH_2)_2$$
 NO_2 NO_2

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For example, a compound of formula (IX) may be reduced by catalytic hydrogenation using a catalyst such as palladium on charcoal to give an amine which may be diazotised using, for example nitrous acid and the product of this reaction may then be reduced using, for example, stannous chloride to give a compound of formula (VII).

According to another general process (D), a compound of formula (I) may be prepared by reduction of a 25 compound of formula (X)

$$R_1R_2NSO_2-CH=CH-1$$

$$R_3$$

$$R_3$$

$$(X)$$

The reduction may be effected using similar reaction conditions to those described for general process (A) above.

Compounds of formula (X) are novel and form a further feature of the invention.

Compounds of formula (X) may be prepared by condensing a compound of formula (XI)

(wherein X represents a leaving atom or group such as a halogen atom for example a bromine atom) with an alkene $R_1R_2NSO_2CH=CH_2$.

The reaction will generally be effected in the presence of a palladium catalyst and a base. The catalyst may be, for example, palladium on charcoal or a palladium salt. Palladium salts which may be employed as catalysts include salts of organic acids such as acetates or salts of inorganic acids such as chlorides or bromides. The base may be, for example, a tertiary nitrogen base such as triethylamine or trin-butylamine or an alkali metal carbonate such as socium carbonate. The reaction may optionally be carried out in the presence of a phosphine, for example a triarylphosphine such as triphenylphosphine or tri-o-tolylphosphine. A phosphine should be present when the process is effected with a compound of formula (XI) wherein X represents a formine atom.

General process (D) may be effected in the presence or absence of solvent. An anhydrous or aqueous reaction medium comprising one or more solvents may be employed. Suitable solvents include nitriles, for example, acetonitrile, alcohols, for example methanol, amides, for example dimethyliomamide, N-methyl-pyrrolidine or hexamethylphosphoramide; and water. The reaction may conveniently be carried out at a temperature of from 25 to 200 °C, preferably 75 to 150° C.

Compounds of formula (XI) may be prepared from known compounds by methods analogous to those to described herein.

According to another general process (E) a compound of formula (I) according to the invention may be converted into another compound of the invention using conventional procedures.

According to one embodiment of general process (E), a compound of general formula (I) wherein one or more of R₁, R₂ and R₄ represent hydrogen atoms may be alkylated using conventional techniques. The reaction may be effected using a suitable alkylating agent such as an alkyl halide, alkyl tosylate or dialkylsulphate. The alkylation reaction may conveniently be carried out in an inert organic solvent such as an amide (e.g. dimethylormamide) or an ether (e.g. tertalydrofuran) represence) at base. Suitable bases include, for example, alkali metal hydrides, such as sodium hydride, alkali metal carbonates, such as sodium carbonate or alkali metal alkoxides such as sodium or potassium methoxide, ethoxide or t-butoxide. The alkylation reaction is conveniently carried out at a temperature of from 25 to 100°C.

According to another general process (F), a compound of formula (I) where R₂ represents a C₃₋₆ alkyl group may be prepared by reduction of the corresponding compound (I) wherein R₂ represents a C₃₋₆ alkenyl group. The reduction process may be effected using the conditions as described above for reduction of the group CH=CH₂ in compounds of formula (II). Compound analogous to compounds of 26 formula (I) but in which R₂ represents a C₃₋₆ alkenyl group may be prepared by methods analogous to those described therein for the preparation of compounds of formula (I).

According to another general process (G), a compound of formula (I) according to the invention, or a salt thereof may be prepared by subjecting a protected derivative of formula (I) or a salt thereof to reaction to remove the protecting group or groups.

Thus, at an earlier stage in the preparation of a compound of formula (I) or a salt thereof it may have been necessary and/or desirable to protect one or more sensitive groups in the molecule to prevent undesirable side reactions.

The protecting groups used in the preparation of compounds of formula (I) may be used in conventional manner. See for example "Protective Groups in Organic Chemistry" Ed.J.F.W. McOmie (Plenum Press 1973) or "Protective Groups in Organic Synthesis" by Theodora W Greene (John Wilev and Sons 1981).

In compounds of formula (I) wherein R₄ represents hydrogen the group NR₄ may be protected for example by protonation or with a conventional amino protecting group. Such groups may include for example arallyl groups, such as benzyl, dipheny/methyl or tripheny/methyl groups; and acyl groups such as N-benzyloxycarbonyl or t-butoxycarbonyl. The indole nitrogen may also be protected, for example by an 40 arallyl group such as benzyl. Thus, compounds of general formula (I) wherein one or more of the groups R₂ and R₃ represent hydrogen may be prepared by deprotection of a corresponding protected compound.

Removal of any amino protecting groups present may be achieved by conventional procedures. Thus an aralkyl group such as benzyl, may be cleaved by hydrogenolysis in the presence of a catalyst (e.g. palladium on charcoal); an acyl group such as N-benzyloxycarbonyl may be removed by hydrolysis with, for example, hydrogen bromide in acetic acid or by reduction, for example by catalytic hydrogenation.

As will be appreciated, in some of the general processes (A) to (F) described above it may be necessary or desired to protect any sensitive groups in the molecule as just described. Thus, a reaction step involving deprotection of a protected derivative of general formula (I) or a salt thereof may be carried out subsequent to any of the above described processes (A) to (F).

Thus, according to a further aspect of the invention, the following reactions may, if necessary and/or desired be carried out in any appropriate sequence subsequent to any of the processes (A) to (F).

(i) removal of any protecting groups; and

(ii) conversion of a compound of formula (I) or a salt thereof into a pharmaceutically acceptable salt or solvate (for example, hydrate) thereof.

Where it is desired to isolate a compound of the invention as a salt, for example as an acid addition salt, this may be achieved by freating the free base of general formula (i) with an appropriate acid, preferably with an equivalent amount, or with creatinine sulphate in a suitable solvent (e.g. aqueous ethanol).

As well as being employed as the last main step in the preparative sequence, the general methods indicated above for the preparation of the compound of the invention may also be used for the introduction of the desired groups at an intermediate stage in the preparation of the required compound. It should therefore be appreciated that in such multi-stage processes, the sequence of reactions should be chosen in 5 order that the reaction conditions do not affect groups present in the molecule which are desired in the final conduct.

The invention is further illustrated by the following Examples. All temperatures are in *C.

Intermediate 1

N-Methyl-3-(1,2,3,6-tetrahydro-l-methyl-4-pyridinyl)-lH-indole-5ethane sulphonamide oxelate

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A solution of N-methyl-H-indole-5-ethanosulphonamide (1.0p) in methanol (50mL) containing potassium hydroxide (5.6g) and N-methyl-4-piperidone (1.0mL) was heated at reflux for 24h, cooled, and the resulting solid filtered off (1.0g). A sample of the solid (0.2g) was dissolved in a hot methanolic solution of 20 oxalic acid (0.0gg), the solution cooled, and the salt precipitated by adding ethyl acetate (20mL) and dry ether (50mL). The salt was filtered off, and dried in vacuo to give the title compound as a solid (0.12g) m.p. 87 *80 * (shrinks) Analysis Found: C,52.2H.5.5N.9.5. C: 1+b:3.N.9.5. C-b:H.0.0 BbO P requires (2.5.2.5H.6.5N.9.7%.

25 Intermediate 2

5-Bromo-3-(1-methyl-4-piperidinyl)-lH-indole

A mixture of 5-bromoindole (39.2g), N-methyl-4-piperidone (25.0g) and potassium hydroxide pollets (12.0g) in methanol 8250ml) was stirred and heated at reflux for 17h then cooled to 5°, with stirring. The mixture was filtered. The residue was washed consecutively with methanol, water, methanol again and other and dried in vacuo to give the intermediate tetrahydropyridine (43.3g) as a powder, with m.p. 256-281° which was used without urther characterisation in the noxt stage, A solution of ethanolic hydrogen cholinds was prepared by adding acetyl chloride (20ml) to ice-cooled, stirred ethanol (1.31). The intermediate tetrahydropyridine (43.2g) was dissolved in a portion (0.951) of this solution. The hydrochloride sall of the intermediate precipitated out. In order to redissolve this salt the suspension was heated on a steam abortions of 2N hydrochlorid acid (10ml), water (15ml) and con. (11N) hydrochlorid acid (10ml) overbon (7.0g) in ethanolic (HCI (0.351 of the above solution) and the mixture was hydrogenated at room temperature and atmospheric pressure until uptake of hydrogen ceased. The mixture was filtered and the solvent was evaporated. The residue was suspended in ethyl acetate (600ml). Sodium carbonate (2N; 30ml) was added, with stirring and the mixture was filtered and the solvent was evaporated. The residue was suspended in ethyl acetate (600ml). Sodium carbonate (2N; 30ml) was added, with stirring and the mixture was filtered. The residue was washed with water and ethyl acetate and dried in vacuo to give the title compound (33.4g) as a powder, mp. 160-165*.

Intermediate 3

5-Bromo-3-[1,2,3,6-tetrahydro-1-(phenylmethyl)-4-pyridinyl]-1H-indole

Freshly distilled 1-Benzyl-4-piperidone (11-7g) was added to a stirred solution of 5-bromoindole (11-0g) in 2M potassium hydroxide in methanol (81m1). The mixture was stirred at reflux for 8h and then allowed to cool to 25° over 8h. The solid was collected by filtration, washed with a mixture of methanol-water (2:1, 2:15mt) and dried in vacuo at 50° for 18h to give the title compound as a crystalline solid (18.6g) m.p. 51 173-175° (docomp).

Intermediate 4

5-Bromo-3-[1-(phenylmethyl)-4-piperidinyl]-lH-indole

A solution of Intermediate 3 (4.00g) in ethanolic hydrogen chloride (330m.t; prepared by the addition of acelyt chloride (1.65g) to ethanol (250m.t) with stirring) was hydrogenated over 5% platinum on carbon (3.0g) at room temperature and atmospheric pressure until hydrogenation was complete. The catalyst was rore removed by filtration. The solid was washed with ethanol (15mt) and the combined filtrate evaporated to give an oily residue. The residue was partitioned between 2M aqueous sodium carbonate (75mt) and ethyl acetate (175mt), the phases separated and the aqueous layer re-extracted with ethyl acetate (100mt). The combined organic layers were then weshed with water (56mt) extracted with saturated brine (56mt), dried (MgSQ₁) and the solvent evaporated to give the title compound as an oil (3.3g). T.Lc. SiO₂ 15 CH₂O₃-E(HD-0.88 NH₁ (100-81) R10-41.

Example 1

N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethansulphonamide

Intermediate 1 (as the free base) (0.36g, 0.001mol) in absolute alcohol (70m.t) and anhydrous dimethyllormamide (5m.t) was hydrogenated, in the presence of 5% palladium on activated carbon (0.36g) at ambient temperature and atmospheric pressure. After 20h, hydrogen absorption 25cm³, theoretical = 24cm³) ceased. The catalyst was filtered off and the solvent removed in vacuo to give an opaque gum which solidified as a soft white solid (0.3g). Purification by flash chromatography (Sorbsi C60 silica gel, CH₂Ct₂/EI0H0.88 ammonia; 50:80:1) gave a colourless oil (0.21g) that was triturated with ether to give the title compound (0.17g) m.p. 156-158*. T.l.c. SiO₂ (CH₂Ct₂/EIOH0.88 ammonia; 50:8:1) Rf 0.4; detection, IU.v., IPA.

Water assay	Found:	0.12% w/w = 0.02mol equiv.		
Analysis	Found : requires	C,60.5;	H,7.3;	N,12.1.
C ₁₇ H ₂₅ N ₃ O ₂ S.O.O2H ₂ O		C,60.8;	H,7.5;	N,12.5%

Example 2

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N-Methyl-3-(l-methyl-4-piperidinyl)-lH-indole-5-ethanesulphonamide

(i) (E)-N-Methyl-2-[3-(1-methyl-4-piperidinyl)-1H-indol-5-yl]ethenesulphonamide

A mixture of Intermediate 2 (1.00g) N-methylethenesulphonamide (530mg), tri-o-tolylphosphine (300mg), palladium acetate (59mg) and triethylamine (730mg) in dry acetonitrile (added to give a total volume of 10m1) was stirred and heated in a sealed vessel at 120° for 125h and then 80° for 15h. The reaction was repeated on the same scale 10 times. In each case the sealed vessel was heated at 100-110° or 3.5h. The sealed vessel were cooled, the contents were combined and the solvent was evaporated. The residue was chromatographed on silica (450g), using a mixture of dichloromethane, ethanol and ammonia (initially 80.81, gradually increasing the polarity to 65:81). The fractions containing the product were combined and evaporated to give a semi-solid. The material was briefly triturated in a mixture of cyclohexane and ethyl acetate (11; 100m1) to give a solid which was filtered and dried to give the title compound (48.50g) as a powder, mp. 190-192°.

(ii) N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide

A solution of the product of stage (i) (5.78g) in a mixture of ethanolic hydrogen chloride [prepared by adding acetyl chloride (1.71g, 21.8mmol) to IMS ethanol (400mt) with stirring] and dimethylformamide

(300mt; added to the above to dissolve the starting material) was hydrogenated at room temperature and atmospheric pressure, using 10% palladium on carbon (5.00g, 50% w/w with water) as the catalyst until uptake of hydrogen ceased. The mixture was filtered and the filtrate was ovaporated to give a solid. The solid was partitioned between 2N sodium carbonate (60mt) and othyl acotate (200m1) and the mixture was 5 heated until the solid had dissolved. The phases were separated, the aqueous phase was extracted with ethyl acetate (200m1) and the combined organic phases were washed with saturated brine (100mt), dried (NasSQ) and evaporated to give a gum. The gum was crystallised from ethyl acetate (60mt) to give the title compound (4.30g) as crystals, with mp. 170-171*

10	Analysis C ₁₇ H ₂₅ N ₃ O ₂ S	Found:	C,60.9;	H,7.6;	N,12.4.
	C ₁₇ H ₂₅ N ₃ O ₂ S	requires	C,60.9;	H,7.5;	N,12.5%.

15 Example 3

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N-Methyl-3-(1-methyl-4-piperidinyl)-lH-indole-5-ethanesulphonamide

A solution of 4-hydrazino-N-methyl-benzenethanesulphonamide (0.5g) and 1-methyl-4-piperidineacetal-dehyde (0.35g) in a mixture of water (10mt) of 2N hydrochloric acid (1.0mt, 2.00mmol) was stirred for 2 days at room temperature. A further quantity of the aldehyde (0.35g) was added and stirring continued for a further 30min. The solution was then basified with 8% sodium bicarbonate to pH8 and extracted with chloroform (3x56mt). The combined organic extracts were dried (Nag-Xol) and evaporated in vacuo to give the crude hydrazone as an oil (1.0g). A solution of the hydrazone (1.0g) in chloroform (20ml) containing polyphosphate seter (10g) was heated at reflux for 8 min. The solution was poured onto ice (200g), stirred for 2h treated with 2M sodium carbonate (20ml) and extracted with chloroform (3 x 50ml).

The combined organic extracts were dried (Na:SO₄), evaporated in vacuo and the residue purified by silash chromatography (silica 9385, 100g) eluting with CH₂Cb/ECH/NH₄ [75:8:1) to give impure material as a yellow oil. Further flash chromatography (silica 9385, 100g) eluting with CH₂Cb/ECH/NH₄ (100:8:1) gave the product as an oil (0.05g). This was crystallised from ethyl acetate to give the title compound solid m.p. 156:157.

T.I.c. SiO₂, CH₂Cl₂/EtOH/NH₃ (50:8:1) Rf 0.6

Example 4

N,N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide

Sodium hydride (80% w/w with paraffin; 124mg) was added cautiously to a stirred solution the product of Example 1 in dry dimethylformamide (20mt). The resultant mixture was stirred at room temperature under nitrogen for 0.25h then a solution of methyl iodide (440mg) in dry dimethylformamide (1mt) was added in a stream. The mixture was stirred at room temperature for 2.5h. The reaction mixture was quenched with water (3mt), evaporated in vacue and the residue was chromatographed on silica (150g), eluting with dichloromethane, ethanol and ammonia (80:10:1) to give a gum. The gum was briefly triturated in diethyl ether and the title compound crystallised out as a powder (238mg), m.p. 170-172*.
T.L.c. SiQ. C(H-Sc):EVBHNH, 50:81), Riv 0.57.

Example 5

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(i) (E)-N,N-Dimethyl-2-[3-(1-methyl-4-piperidinyl)-lH-indol-5-yl]ethenesulphonamide

A mixture of 5-Bromo-3-(1-methyl-4-piperidinyl)-1H-indole (2.0g) N.N-dimethylethenesulphonamide (1.184g), tri-o-tolylphosphine (0.6g) palladium acetate ($\overline{O.1}$ g), triethylamine (1.0m1) and anhydrous acetoni-

EP 0 303 507 B1

trile (12m1) was heated in two 10m1 sealed vessels, with stirring at 107° (oil bath temp) for 2.25h. The reaction mixtures were combined, the solvent removed by rotary evaporation and the residual foam purified by flash chromatography eluting with dichloromethane/ethane/10.88 ammonia (100.8:1). Rotary evaporation of the appropriate fractions gave the product as a foam (1920)

5 T.I.c. SiO₂ isopropanol/ethanol/water/0.88 ammonia (20:20:8:1) Rf 0.5 (major) + 0.55 (minor) + 0.4 (trace).

(ii) N,N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide

A solution of the product of stage (1) (1.5g) in ethanol (200m1) was added to a slurry of 5% palladium on activated carbon (1.5g) in ethanol (100m1). The resulting mixture was hydrogenated at 65psi at room temperature for 17h. The mixture was filtered and the filtrate evaporated to leave a solid (1.0g) which was washed with isopropanol (3x20m1) to give a solid (0.8g) m.p. 215-225*

Crystallisation from hot ethanol (60mt) gave the title compound as microneedles (0.29g) m.p. 228-232* T.I.c. SiO₂ isopropanol/ether/water/0.88 ammonia (20:20:8:1) Rf 0.5.

Example 6

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3-(1-Methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide

(i) (E)-2-[3-(1-methyl-4-piperidinyl)-1H-indol-5-yl]

ethenesulphonamide

A mixture of Intermediate 2 (2.0g), vinyl sulphonamide (0.88g) palladium acetate (100mg), triin(t-loty)-phosphine (0.8g), triethyalmine (1.0mt), and acetonitrie (14mt) was separated into two equal portions and so placed into two sealed vessels (10mt) and heated at 100° for 4h. A further quantity of the vinyl sulphonamide (0.22g) was added to each sealed vessel and the mixture was heated at 100° for a further 16h. The resulting mixture was evaporated to dryness in vacuo and the residue purified by flash chromatography (sliica 9385, 400g) eluting with CHCl₂/EtOH/NH₃ (100:8:1 to 75:8:1) to give the title compound as a solid (0.8 m.p. 208-200°.

(ii) 3-(1-methyl-4-piperidinyl)-lH-indole-5-ethanesulphonamide

A mixture of the product of stage (i) (0.8g) in ethanolic hydrogen chloride (80m1.) was hydrogenated over pre-reduced 10% palladium on carbon (50% paste with water, 0.8g) until uptake ceased. The catalyst was filtered off, washed with hot ethanol (50m1) and the filtrate evaporated in vacuo to give crude material (0.15g). The catalyst residues were then warmed (70·1) with 2N hydrochloric acid (200m1), filtered and the filtrate evaporated to dryness in vacuo (acertoped with toluene). The residue was combined with the crude product obtained above and purified by flash chromatography (silica 9385, 100g) eluting with 45 CH₂CH₂CHOHNH₃ (50.8:1) to give the title compound as a solid (0.2g) m.p. >95° (floams). T.L.C. SiQ. C.H.C.E/CEDHNH₃ (25.8:1) flif 0.5.

Example 7

50 N-Methyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide hydrochloride

(i) (E)-N-Methyl-2-[3-[1-(phenylmethyl)-4-piperidinyl]-1H-indol-5-yl]ethenesulphonamide

In each of three sealed vessels, a mixture of Intermediate 4 (1.10g), N-methyl ethenesulphonamide (422mg), triethylamine (843u.1) tri-o-tolyphosphine (242mg) and palladium acetate (39mg) in dry acetonitritic (volume made up to 10m1) was stirred and heated at 100° for 4h. After cooling to 25° the contents of the vessels were combined and the solvent evaporated in vacuo at 40° to give an oily residue. This residue was purified by column chromatography on silica gel (Merck 722, 300g) eluting with a mixture of the content of the

dichloromethane:ethanoi:0.88 ammonia (300:8:1 to 200:8:1). The appropriate fractions were combined, and the solvent evaporated in vacuo to give the title-compound a foam (2.14g). T.L.c. SiO-Gio-Gio-Ei-TOH-0.88NH, (200:8:11) RT 0.41.

5 (ii) N-Methyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide hydrochloride

A solution of the product of stage (i) (2.14g) in ethanolic hydrogen chloride (350m.t, prepared by the addition of acetyl chloride (850mg) in ethanol (350m1) with stirringl was hydrogenated over pre-reduced 10% palladium on charcoal (6.4g) at 25° and 1 atmosphere pressure for t8h. The reaction mixture was purged with nitrogen and a solution of ammonium formate (8.2g) in methanol (100m1) added. The mixture was stirred and brought to reflux under nitrogen for 10min, cooled at 25° and the catalyst removed by filtration. Evaporation of the filtrate in vacuo gave a solid residue (8.5g) which was redissolved in water (75m1) and saturated with solid sodium chloride. The resultant precipitate was collected by filtration, washed with ice-cold water (1.5m1) and ether (10mt) and dried in vacuo at 45° for 18h to give the title sompound as a crystalline solid (640mg) mp. 253-255°.

T.I.c. SiO₂ CH₂Cl₂:EtOH:0.88NH₃ (25:8:1), Rf 0.14.

Example 8

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N-Ethyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide

(i) (E)-N-Ethyl-2-[3-[1-(phenylmethyl)-4-piperidinyl]-1H-indol-5-yl]ethene sulphonamide

Into each of two 10mt sealed vessels were placed palladium acetate (50mg), trit-otolyphosphine (300mg), triethylamine (650mg), N-ethylethenesulphonamide (275mg) and Intermediate 4 (710mg). Each mixture was made up to 10mt with dry acetonitrile. The vessels were heated at 100° for 16h then left at room temperature for 4 days. The contents of the sealed vessels were combined and the solvent and triethylamine were removed in vacuo. The residue was chromatographed on silica (205mg; Merck 9385), eluting with dichloromethane, ethanol and ammonia (100:8:1) as the eluant, to give a foam (759mg). The foam was crystallised from a hot mixture of ethyl acetate and cyclohexane to give the title compound - (582mn) as microcrystals no., 178-180°.

(ii) N-Ethyl-3-(4-piperidinyl)-lH-indole-5-ethanesulphonemide

A solution of the product of stage (i) (370mg) in ethanolic hydrogen chloride [prepared by adding acetyl chloride (105mg, 1.34mmol) to IMS ethanol (50m1), with stirring] was hydrogenated over pre-reduced 10% palladium oxide on carbon (50% w/w with H₂O; 1.13g), at room temperature and atmospheric pressure until uptake of hydrogen ceased. The mixture was filtered and the filtrate was evaporated to give a foam (280mg) which was dissolved in methanol (4m1). Sodium carbonate (2N): 2m1) was added and the solvent was exporated. The residue was partitioned between water (10m1) and ethyl acetate (50m1). The aqueous phase was extracted with ethyl acetate (50m1) and the combined organics were dried (Na₂SO₄) and evaporated to give a gum 235mg) which was crystallised from a mixture of ethyl acetate and ether (10m1; mainly ethyl acetate) to give the title compound (104mg) as a powder m.p. 95-100 *.
T.L. SiQC (2H-GL:ELFOHNH)_E (258:1), RIO 3.

50 Example 9

N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide hydrochloride

A solution of the product of Example 1 (50mg) in hot ethanol (0.5mt) was added to ethanolic hydrogen chloride [prepared by adding acetyl chloride (33mg, 0.420mmol) to ethanol (1mt) at room temperature] in a stream with stirring, at room temperature. A solid crystallised out from the initially clear solution. The suspension was stirred and cooled to 5 * over 15min then filtered under suction. The residue was washed with a little ethanol and then dried at 60 * in vacuo for 1th to give the title compound (44mg) as

microcrystals, m.p. 237-239°.

T.I.c. SiO₂ (CH₂Cl₂:EtOH:NH₃ 50:8:1). Rf 0.45.

The following examples illustrate pharmaceutical formulations according to the invention containing N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethansulphonamide as the active ingredient. Other compounds of the invention may be formulated in a similar manner.

TABLETS FOR ORAL ADMINISTRATION

A. Direct Compression

1.

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	mg/tablet
Active ingredient Magnesium Stearate BP	49 0.65
Anhydrous Lactose	81

The active ingredient is sieved and blended with the anhydrous lactose and magnesium stearate. The resultant mix is compressed into tablets using a Manesty F3 tablet machine fitted with 8.0mm concave punches.

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	mg/tablet
Active ingredient	49
Magnesium Stearate BP	0.7
Microcrystalline Cellulose NF	91

The active ingredient is sieved and blended with the microcrystalline cellulose and magnesium stearate. The resultant mix is compressed into tablets using a Manesty F3 tablet machine fitted with 8.0mm concave punches.

35 B WET GRANULATION

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	mg/tablet
Active ingredient	7.0
Lactose BP	146.5
Starch BP	30.0
Pregelatinised Maize Starch BP	15.0
Magnesium Stearate BP	1.5
Compression weight	200.0

The active ingredient is sieved through a suitable sieve and blended with lactose, starch and progelatinised maize starch. Suitable volumes of purified water are added and the powders are granulated. After drying, the granules are screened and blended with the magnesium stearate. The granules are then compressed into tablets using suitable diameter punches.

Tablets of other strengths may be prepared by altering the ratio of active ingredient to lactose or the compression weight and using punches to suit.

The tablets may be film coated with suitable film-forming materials, such as hydroxypropyl methylcellulose, using standard techniques. Alternatively the tablets may be sugar coated, or enteric coated.

CAPSULES

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	mg/capsule
Active ingredient	49.00
*Starch 1500	150.00
Magnesium Stearate BP	1.00
Fill Weight	200.00

^{*} A form of directly compressible starch,

The active ingredient is sieved and blended with the excipients. The mix is filled into size No.2 hard gelatin capsules using suitable machinery. Other doses may be prepared by altering the fill weight and if necessary changing the capsule size to suit.

SYRUP

20	Sucrose Free Presentation	mg/5ml dose
	Active Ingredient	49.00
25	Hydroxypropylmethylcellulose USP (viscosity type 4000)	22.5
30	Buffer) Flavour) Colour) Preservative) Sweetener)	as required
	Purified Water BP to	5.0ml

The hydroxypropylmethylcollulose is dispersed in hot water, cooled and then mixed with an aqueous solution containing the active ingredient and the other components of the formulation. The resultant solution is adjusted to volume and mixed. The syrup is clarified by filtration.

SUSPENSION

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45		mg/5ml dose
	Active ingredient	49.00
	Aluminium monostearate	75.00
50	Sweetening agent) Flavour) Colour)	as required
	Fractionated coconut oil to	5.00ml

The aluminium monostearate is dispersed in about 90% of the fractionated coconut oil. The resulting suspension is heated to 115°C while stirring and then cooled. The sweetening agent, flavour and colour are added and the active ingredient is suitably dispersed. The suspension is made up to volume with the remaining fractionated coconut oil and mixed.

SUB-LINGUAL TABLET

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mg/tablet
Active Ingredient 49.00
Compressible Sugar NF 50.5
Magnesium Stearate BP 0.5
Compression Weight 100.0

The active ingredient is sieved through a suitable sieve, blended with the excipients and compressed using suitable punches. Tablets of other strengths may be prepared by altering either the ratio of active ingredient to excipients or the compression weight and using punches to suit.

SUPPOSITORY FOR RECTAL ADMINISTRATION

Active ingredient	49.0mg
* Witepsol H15 to	1.0g

^{*} A proprietary grade of Adeps Solidus Ph. Eur.

A suspension of the active ingredient in molten Witepsol is prepared and filled, using suitable machinery, into 1g size suppository moulds.

INJECTION FOR INTRAVENOUS ADMINISTRATION

	mg/ml
Active Ingredient Sodium Chloride Intravenous Infusion, BP, 0.9% w/v Batch Size	0.896 to 1 ml 2500ml

The active ingredient is dissolved in a portion of the Sodium Chloride Intravenous Infusion, and the solution made to volume with the Sodium Chloride Intravenous Infusion, and the solution thoroughly mixed. The solution is filled into clear, Type I, glass 10ml ampoules and sealed under a nitrogen headspace by fusion of the glass. The ampoules are sterilised by autoclaving at 121 °C for not less than 15 minutes.

FOR INHALATION

Inhalation Cartridges	
	mg/cartridge
Active ingredient (micronised) Lactose BP	0.56 25.00

The active ingredient is micronised in a fluid energy mill to a fine particle size range prior to blending with normal tabletting grade lactose in a high energy mixer. The powder blend is filled into No. 3 hard gelatin capsules on a suitable encapsulating machine. The contents of the cartridges are administered using a powder inhaler such as the Glaxo Rotahaler.

Metered Dose Pressurised Aerosol		
Suspension Aerosol	mg/metered dose	Per can
Active ingredient (micronised) Oleic Acid BP	0.280 0.020	73.92mg 5.28mg
Trichlorofluoromethane BP Dichlorodifluoromethane BP	23.64 61.25	5.67g 14.70g

The active ingredient is micronised in a fluid energy mill to a fine particle size range. The oleic acid is mixed with the trichloromethane at a temperature of 10-15°C and the micronised drug is mixed into the solution with a high shear mixer. The suspension is metered into aluminium aerosol cans and suitable metering valves, delivering 85mg of suspension are crimped onto the cans and the dichlorodifluoromethane is pressure filled into the cans through the valves.

Nasal Spray		
	% w/v	
Active Ingredient	7.0	
Preservative	as required	
Sodium Chloride BP		
Purified Water BP to	100	
Shot Weight	100mg (equivalent to 7mg active ingredient)	

The active ingredient, preservative and sodium chloride are dissolved in a portion of the water, the solution made to volume with the water and the solution thoroughly mixed.

The pH may be adjusted, using acid or alkali, to that of optimum stability and/or to facilitate solution of the active ingredient. Alternatively, suitable buffer salts may be used.

Claims

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Claims for the following Contracting States: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

1. A compound of formula (I)

$$R_1R_2NSO_2(CH_2)_2$$

whorein

R₁ represents a hydrogen atom or a C_{1−6} alkyl group;

R₂ represents a hydrogen atom or a C₁₋₆ alkyl group;

R₃ represents a hydrogen atom

R₁ represents a hydrogen atom or a C₁₋₃ alkyl group;

and pharmaceutically acceptable salts and solvates thereof.

- 2. A compound according to Claim 1 wherein R₁ represents a hydrogen atom or a C₁₋₃ alkyl group.
- 55 3. A compound according to Claim 1 or 2 wherein R2 represents a hydrogen atom or a C1-3 alkyl group.
 - A compound according to any of Claims 1 to 3 wherein R₂ represents a C₁-₃ alkyl group.

- A compound according to any of Claims 1 to 4 wherein R_t represents a C₁₋₃ alkyl group.
- N-methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- N,N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- N-Ethyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
 - N-Methyl-3-(4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- 15 10. 3-(1-Methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
 - 11. A compound according to any of Claims 1 to 10 for use in therapy.

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- 20 12. The use of a compound according to any of Claims 1 to 10 in the preparation of a medicament for use in the treatment of conditions associated with cephalic pain.
 - 13. The use of a compound according to any of Claims 1 to 10 in the preparation of a medicament for use in the treatment of migraine cluster headache, chronic paroxysmal hemicrania or headache associated with vascular disorders and in alleviation the symptoms associated therewith.
 - 14. A pharmaceutical composition which comprises at least one compound of formula (I) as defined in Claim 1 or a pharmaceutically acceptable saitor solvate thereof together with one or more pharmaceutically acceptable carriers or excipients.
 - 15. A pharmaceutical composition as claimed in Claim 14 adapted for oral, parenteral or intranasal administration.
- 16. A pharmaceutical composition according to Claim 14 or 15 which is formulated in unit dosage form comprising 0.1mg to 100mg of active ingredient.
 - 17. A process for the preparation of a compound of formula (I) as defined in Claim 1 or a pharmaceutically acceptable salt or solvate thereof which comprises (A) reducing a compound of formula (II)

 $R_1R_2NSO_2(CH_2)_2 - \sqrt[4]{NR_4}$ (II)

wherein R1, R2, R3 and R4 are as defined in Claim 1.

(B) condensing an amine of formula $R_1\,R_2\,NH$ (wherein R_1 and R_2 are as defined in Claim 1) with a compound of formula (V)

$$HOSO_2(CH_2)_2 \xrightarrow{\uparrow} NR_4$$

$$R_3$$

$$(V)$$

(wherein R_0 and R_4 are as defined in Claim 1), or an acylating agent corresponding thereto or a salt or protected derivative thereof; or

(C) cyclising a compound of formula (VI)

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$${\sf R_1R_2NSO_2(CH_2)_2} = \underbrace{{\color{red} {\color{blue} {\color{b} {\color{blue} {\color{b} {\color{blue} {\color{blue} {\color{blue} {\color{blue} {\color{blue} {\color{b} {\color{blue} {\color{b} {\color{b} {\color{blue} {\color{b} {$$

wherein R_1 , R_2 , R_3 and R_4 are as defined in Claim 1;

(D) reducing a compound of formula (X)

$$\mathsf{R_1R_2NSO_2CH=CH-} \bigvee_{\mathsf{R_3}}^{\mathsf{NR_4}} \mathsf{NR_4} \tag{X}$$

(wherein R₁, R₂, R₃ and R₄ are as defined in Claim 1).

- (E) subjecting another compound of formula (I) to an interconversion reaction,
- (F) in order to prepare a compound of formula (I) in which R_2 represents a C_{3-6} alkyl group reducing the corresponding compound in which R_2 represents a C_{3-6} alkenyl group;
- (G) subjecting a protected derivative of a compound of formula (f) or a salt thereof to reaction to remove the protecting group or groups; and if necessary or desired subjecting the compound resulting from any of steps (A) to (F) to one or two further reactions comprising:

i) removing any protecting groups

ii) converting a compound of formula (I) or a salt thereof into a pharmaceutically acceptable salt of solvate thereof.

Claims for the following Contracting States: GR, ES

1. A process for the preparation of a compound of formula (I)

$$R_{1}R_{2}NSO_{2}(CH_{2})_{2}$$

wherein

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R₁ represents a hydrogen atom or a C₁₋₆ alkyl group;

R2 represents a hydrogen atom or a C1-6 alkyl group;

R₃ represents a hydrogen atom;

R4 represents a hydrogen atom or a C1-3 alkyl group;

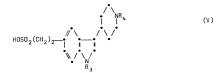
and pharmaceutically acceptable salts and solvates thereof which comprises :

(A) reducing a compound of formula (II)

$$R_1R_2NSO_2(CH_2)_2$$

(wherein R₁, R₂, R₃ and R₄ are as defined in Claim 1);

(B) condensing an amine of formula $R_1\,R_2\,NH$ (wherein R_1 and R_2 are as defined in Claim 1) with a compound of formula (V)



(wherein R_2 and R_4 are as defined in Claim 1), or an acylating agent corresponding thereto or a salt or protected derivative thereof; or

(C) cyclising a compound of formula (VI)

(wherein R₁, R₂, R₃ and R₄ are as defined in Claim 1);

(D) reducing a compound of formula (X)

$$\mathbf{R_1}\mathbf{R_2} \mathsf{NSO}_2 \mathsf{CH} = \mathsf{CH} - \bigvee_{\mathbf{R_3}} \bigvee_{\mathbf{R_3}} \mathsf{NSO}_2 \mathsf{CH} = \mathsf{CH} - \bigvee_{\mathbf{R_3}} \mathsf{NSO}_2 \mathsf{CH} = \mathsf{CH} + \bigvee_{\mathbf{R_3}} \mathsf{CH} + \bigvee_{\mathbf{R_3}} \mathsf{CH} +$$

(wherein R1, R2, R4 and R4 are as defined in Claim 1):

or

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- (E) subjecting another compound of formula (I) to an interconversion reaction;
- (F) in order to prepare a compound of formula (I) in which R_2 represents a C_{3-6} alkyl group reducing the corresponding compound in which R_2 represents a C_{3-6} alkenyl group;
- (G) subjecting a protected derivative of a compound of formula (I) or a salt thereof to reaction to remove the protecting group or groups; and if necessary or desired subjecting the compound resulting from any of stops (A) to (F) to one or two further reactions comprising:
 - i) removing any protecting groups
 - ii) converting a compound of formula (I) or a salt thereof into a pharmaceutically acceptable salt or solvate thereof.
- 2. A process according to Claim 1 wherein R₁ represents a hydrogen atom or a C₁₋₃ alkyl group.
- 3. A process according to Claim 1 or 2 wherein R2 represents a hydrogen atom or a C1-3 alkyl group.
- 45 4. A process according to any of Claims 1 to 3 wherein R₂ represents a C₁-3 alkyl group.
 - 5. A process according to any of Claims 1 to 4 wherein R4 represents a C1-3 alkyl group.
 - A process according to Claim 1 wherein the product is N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indole-5ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
 - A process according to Claim 1 wherein the product is N.N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- 8. A process according to Claim 1 wherein the product is N-Ethyl-3-(4-piperidinyl)-1H-indole-5-ethanesul-phonamide and pharmaceutically acceptable salts and solvates thereof.

- 9. A process according to Claim 1 wherein the product is N-Methyl-3-(4-piperidinyl)-1H-indole-5ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- 10. A process according to Claim 1 wherein the product is 3-(1-Methyl-4-piperidinyl)-1H-indole-5-ethanesulphonamide and pharmaceutically acceptable salts and solvates thereof.
- 11. A process according to any of Claims 1 to 10 wherein in step A or D the reaction is effected in the presence of hydrogen and a catalyst at a temperature of from ~10 to +50 * C.
- 12. A process according to any of Claims 1 to 10 wherein in step (B) the reaction is effected using an acylating agent corresponding to formula (V) at a temperature of from ~70 to +150 °C, optionally in the presence of a base.
- 13. A process according to any of Claims 1 to 10 wherein in step (C) the reaction is effected at a temperature of from 20 to 200 °C.
 - 14. A process according to any of Claims 1 to 10 wherein in step (E) a compound of formula (I) wherein one or more of R1, R2 and R4 are alkyl groups is prepared by alkylating a corresponding compound of formula (1) wherein one or more of R1, R2 and R4 represent hydrogen atoms.
 - 15. A process for preparing a pharmaceutical composition comprising admixing a compound of formula (I) as defined in Claim 1 or a pharmaceutically acceptable salt or solvate thereof together with one or more pharmaceutically acceptable carriers or excipients.

25 Patentansprüche

Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

1. Verbindung der Formel (I):

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$$R_1R_2NSO_2(CH_2)_2$$

$$R_1R_2NSO_2(CH_2)_3$$

$$R_3$$

$$R_3$$

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worin

R₁ für ein Wasserstoffatom oder eine C₁-6-Alkylgruppe steht;

R₂ für ein Wasserstoffatom oder eine C1-6-Alkylgruppe steht;

Ra für ein Wasserstoffatom steht:

R4 für ein Wasserstoffatom oder eine C1-3-Alkylgruppe steht,

- 45 sowie die pharmazeutisch annehmbaren Salze und Solvate davon.
 - 2. Verbindung nach Anspruch 1, dadurch gekennzeichnet, daß R1 für ein Wasserstoffatom oder eine C₁₋₃-Alkylgruppe steht.
 - 3. Verbindung nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß R2 für ein Wasserstoffatom oder eine C1-3-Alkylaruppe steht.
- Verbindung nach einem der Ansprüche 1 bis 3. dadurch gekennzeichnet, daß Re für eine C1-3-55 Alkylgruppe steht.
 - Verbindung nach einem der Ansprüche 1 bis 4, dadurch gekennzeichnet, daß R₄ für eine C₁₋₃-Alkylaruppe steht.

- N-Methyl-3-(1-methyl-4-piperidinyl)-1H-indol-5-ethansulfonamid und die pharmazeutisch annehmbaren Salze und Solvate davon.
- N,N-Dimethyl-3-(1-methyl-4-piperidinyl)-1H-indol-5-ethansulfonamid und die pharmazeutisch annehmbaren Salze und Solvate davon.
- N-Ethyl-3-(4-piperidinyl)-1H-indol-5-ethansulfonamid und die pharmazeutisch annehmbaren Salze und Solvate davon.
- N-Methyl-3-(4-piperidinyl)-1H-indol-5-ethansulfonamid und die pharmazeutisch annehmbaren Salze und Solvate davon.
 - 3-(1-Methyl-4-piperidinyl)-1H-indol-5-ethansulfonamid und die pharmazeutisch annehmbaren Salze und Solvate davon.
 - 11. Verbindung nach einem der Ansprüche 1 bis 10 zur Verwendung in der Therapie.

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- 12. Verwendung einer Verbindung nach einem der Ansprüche 1 bis 10 zur Herstellung eines Arzneimittels zur Verwendung bei der Behandlung von Zuständen im Zusammenhang mit Schmerzen im Kopfbereich.
- 13. Verwendung einer Verbindung nach einem der Ansprüche 1 bis 10 zur Herstellung eines Arzneimittels zur Verwendung bei der Behandlung von Migräne, Cluster Headache, chronischer anfallsartiger Hemikranie oder Kopfschmerzen im Zusammenhang mit vaskulären Störungen und zur Linderung der damit verbundenen Symotome.
- 14. Pharmazeutisches Präparat, dadurch gekennzelchnet, daß es mindestens eine Verbindung der Formel (i) nach Anspruch 1 oder ein pharmazeutisch annehmbares Salz oder Solvat davon zusammen mit einem oder mehreren pharmazeutisch annehmbaren Trägern oder Exzipientien enthält
- 15. Pharmazeutisches Präparat nach Anspruch 14, dadurch gekennzeichnet, daß es für die orale, parenterale oder intranasale Verabreichung angepaßt ist.
- Pharmazeutisches Präparat nach Anspruch 14 oder 15, dadurch gekennzeichnet, daß es in Einheitsdosisform mit 0,1 mg bis 100 mg Wirkstoff formuliert ist.
- 17. Verfahren zur Herstellung einer Verbindung der Formel (I) nach Anspruch 1 oder eines pharmazeutisch annehmbaren Salzes oder Solvats davon, dadurch gekennzelchnet, daß man (A) eine Verbindung der Formel (II);

$$\mathsf{R_1}\mathsf{R_2}\mathsf{NSO}_2(\mathsf{CH}_2)_2 - \bigvee_{\mathsf{R}_1}^{\mathsf{N}_1} \bigvee_{\mathsf{R}_2}^{\mathsf{N}_2} \bigvee_{\mathsf{R}_2}$$

worin R₁, R₂, R₃ und R₄ wie in Anspruch 1 definiert sind, reduziert; oder (B) ein Arnin der Formel R₁ R₂NH (worin R₁ und R₂ wie in Anspruch 1 definiert sind) mit einer Verbindung der Formel (V):

$$HOSO_{2}(CH_{2})_{2} \xrightarrow{\uparrow} \bigvee_{R_{3}} \bigvee_{R_{3}} (V)$$

(worin R₃ und R₄ wie in Anspruch 1 definiert sind) oder einem Acylierungsmittel, das dieser entspricht, oder einem Salz oder einem geschützten Derivat davon kondensiert; oder (C) eine Verbindung der Formel (VI):

$$R_1R_2NSO_2(CH_2)_2$$
 (VI)

worin R₁, R₂, R₃ und R₄ wie in Anspruch 1 definiert sind, cyclisiert; oder (D) eine Verbindung der Formel (X):

$$R_1R_2NSO_2CH=CH$$
 NR_4
 NR_4
 NR_4
 NR_4

(worin R1, R2, R3 und R4 wie in Anspruch 1 definiert sind) reduziert:

- (E) eine andere Verbindung der Formel (I) einer Interkonversionsreaktion unterwirtt; oder (F) zur Herstellung einer Verbindung der Formel (I), bei der R₂ für eine C₃₋₆-Alkylgruppe steht, die entsprechende Verbindung, bei der R₂ für eine C₃₋₅-Alkenvloruppe steht, reduziert; oder
- (G) ein geschütztes Derivat einer Verbindung der Formel (f) oder eines Salzes davon einer Reaktion zur Entfernung der Schutzgruppe oder der Schutzgruppen unterwirft; und erforderlichenfalls oder gewünschenfalls die von einer der Stufen (A) bis (F) resultierende Verbindung einer oder zwei weiteren Reaktionen, umfassend:
 - (i) die Entfernung irgendwelcher Schutzgruppen,
- (ii) die Umwandlung einer Verbindung der Formel (I) oder eines Salzes davon in ein pharmazeutisch annehmbares Salz oder Solvat davon,
- unterwirft.

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Patentansprüche für folgende Vertragsstaaten : GR, ES

1. Verfahren zur Herstellung einer Verbindung der Formel (I):

$$R_{1}R_{2}NSO_{2}(CH_{2})_{2}$$

$$NR_{4}$$

$$NR_{4}$$

$$R_{3}$$

$$R_{3}$$

worin

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R₁ für ein Wasserstoffatom oder eine C₁₋₆-Alkylgruppe steht;

R₂ für ein Wasserstoffatom oder eine C₁₋₆-Alkylgruppe steht;

R₃ für ein Wasserstoffatom steht:

Rt für ein Wasserstoffatom oder eine C1-3-Alkylgruppe steht,

sowie der pharmazeutisch annehmbaren Salze und Solvate davon, dadurch gekennzeichnet, daß man

(A) eine Verbindung der Formel (II):

$$R_1R_2NSO_2(CH_2)_2 - \bigvee_{\substack{i \\ i \\ N_1}} \bigvee_{\substack{i \\ N_3}} (II)$$

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worin R₁, R₂, R₃ und R₄ wie in Anspruch 1 definiert sind, reduziert; oder

(B) ein Amin der Formel R_1R_2NH (worin R_1 und R_2 wie in Anspruch 1 definiert sind) mit einer Verbindung der Formel (V):

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(worin R_3 und R_4 wie in Anspruch 1 definiert sind) oder einem Acylierungsmittel, das dieser entspricht, oder einem Salz oder einem geschützten Derivat davon kondensiert; oder

(C) eine Verbindung der Formel (VI):

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$$R_{1}R_{2}NSO_{2}(CH_{2})_{2} - \bigvee_{NR_{3}N=CHCH_{2}} \bigvee_{NR_{4}} NR_{4}$$
 (VI)

worin R_1 , R_2 , R_3 und R_4 wie in Anspruch 1 definiert sind, cyclisiert; oder (D) eine Verbindung der Formel (X):

$$\mathbf{R_{1}R_{2}NSO_{2}CH=CH} = \underbrace{\mathbf{N}\mathbf{R_{1}}}_{\mathbf{R_{3}}} \underbrace{\mathbf{N}\mathbf{R_{4}}}_{\mathbf{R_{3}}} \tag{X}$$

(worin R1, R2, R3 und R4 wie in Anspruch 1 definiert sind) reduziert;

(E) eine andere Verbindung der Formel (I) einer Interkonversionsreaktion unterwirft; oder

(F) zur Herstellung einer Verbindung der Formel (I), bei der R₂ für eine C₃₋₆-Alkylgruppe steht, die entsprechende Verbindung, bei der R₂ für eine C₃₋₆-Alkenylgruppe steht, reduziert; oder

(G) ein geschütztes Derivät einer Verbindung der Formel (I) oder eines Salzes davon einer Reaktion zur Entfernung der Schutzgruppe oder der Schutzgruppen unterwirft; und erforderlichenfalls oder gewünschtenfalls die von einer der Stufen (A) bis (F) resultierende Verbindung einer oder zwei weiteren Reaktionen, umfassend:

(i) die Entfernung irgendwelcher Schutzgruppen.

 (ii) die Umwandlung einer Verbindung der Formel (I) oder eines Satzes davon in ein pharmazeutisch annehmbares Satz oder Solvat davon, unterwirft.

- Verfahren nach Anspruch 1, dadurch gekenn zeichnet, daß R₁ für ein Wasserstoffatom oder eine C₁₋₃-Alkylgruppe steht.
 - Verfahren nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß R₂ für ein Wasserstoffatom oder eine C₁₋₃-Alkylgruppe steht.
 - Verfahren nach einem der Ansprüche 1 bis 3, dadurch gekennzeichnet, daß R₂ für eine C₁₋₃-Alkylgruppe steht.
 - Verfahren nach einem der Ansprüche 1 bis 4, dadurch gekennzeichnet, daß R₄ für eine C₁₋₃-Alkylgruppe steht.
 - Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Produkt N-Methyl-3-(1-methyl-4piperidinyl)-1H-indol-5-ethansulfonamid und pharmazeutisch annehmbare Salze und Solvate davon ist.
- Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Produkt N,N-Dimethyl-3-(1-methyl-4piperidinyl)-1H-indol-5-ethansulfonamid und pharmazeutisch annehmbare Salze und Solvate davon ist.

- Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Produkt N-Ethyl-3-(4-piperidinyl)-1Hindol-5-ethansulfonamid und pharmazeutisch annehmbare Salze und Solvate davon ist.
- Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Produkt N-Methyl-3-(4-piperidinyl)-1Hindol-5-ethansulfonamid und pharmazeutisch annehmbare Salze und Solvate davon ist.
 - Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Produkt 3-(1-Methyl-4-piperidinyl)-1Hindol-5-ethansulfonamid und pharmazeutisch annehmbare Salze und Solvate davon ist.
- 10 11. Verfahren nach einem der Ansprüche 1 bis 10, dadurch gekennzeichnet, daß man in Stufe (A) oder (D) die Reädkich in Gegenwart von Wasserstoff und eines Katalysators bei einer Temperatur von -10 bis +50 °C vornimmt.
- Verfahren nach einem der Ansprüche 1 bis 10, dadurch gekennzeichnet, daß man in Stufe (B) die Reaktion unter Verwendung eines Acylierungsmittles, das der Formel (V) entsprich, bei einer Temperatur von -70 bis + 150 °C, gegebenenfalls in Gegenwart einer Base, vornimmt.
 - Verfahren nach einem der Ansprüche 1 bis 10, dadurch gekennzelchnet, daß man in Stufe (C) die Reaktion bei einer Temperatur von 20 bis 200 °C vornimmt.
 - 14. Verfahren nach einem der Ansprüche 1 bis 10, dadurch gekennzelchnet, daß man in Stufe (E) eine Verbindung der Formel (I), bei der eine oder mehrere von R₁, R₂ und R₂ Alkylgruppen ist bzw. sind, dadurch herstellt, daß man eine entsprechende Verbindung der Formel (I), bei der eine oder mehrere von R₁, R₂ und R₃ Wasserstoffatome ist bzw. sind, alkyliert.
 - 15. Verfahren zur Herstellung eines pharmazeutischen Präparats, dadurch gekennzeichnet, daß man eine Verbindung der Formel (I) nach Anspruch 1 oder ein pharmazeutisch annehmbrers Batz oder Solvat davon mit einem oder mehreren pharmazeutisch annehmbaren Trägen oder Exzigientien vermischt.

30 Revendications

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Revendications pour les Etats contractants suivants : AT. BE. CH. DE. FR. GB. IT. LI. LV. NL. SE

1. Composé de formule (I).

$$\mathsf{R}_1\mathsf{R}_2\mathsf{NSO}_2(\mathsf{CH}_2)_2 - \bigvee_{\mathsf{R}_3}^{\mathsf{NR}_4} \mathsf{NR}_4 \qquad (\mathsf{I}$$

- dans laquelle
- R₁ représente un atome d'hydrogène ou un groupe alkyle en C₁₋₆ ;
- R₂ représente un atome d'hydrogène ou un groupe alkyle en C₁₋₆;
- R₃ représente un atome d'hydrogène ;
- Ri représente un atome d'hydrogène ou un groupe alkyle en C₁₋₃ :
- et ses sels et produits de solvatation pharmaceutiquement acceptables.
- Composé selon la revendication 1, dans lequel R₁ est un atome d'hydrogène ou un groupe alkyle en C₁₋₃.
- Composé selon la revendication 1 ou 2, dans lequel R₂ est un atome d'hydrogène ou un groupe alkyle en C₁₋₃.

- Composé selon l'une quelconque des revendications 1 à 3, dans lequel R₂ est un groupe alkyle en C₁₋₃.
- Composé selon l'une quelconque des revendications 1 à 4, dans lequel R₄ est un groupe alkyle en C₁₋₃.
 - N-méthyl-3-(1-méthyl-4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
- N,N-diméthyl-3-(1-méthyl-4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
 - N-éthyl-3-(4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
 - N-méthyl-3-(4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
- 10. 3-(1-mèthyl-4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmace ceutiquement acceptables.
 - 11. Composé selon l'une quelconque des revendications 1 à 10, pour une utilisation thérapeutique.

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- 12. Utilisation d'un composé selon l'une quelconque des revendications 1 à 10, pour préparer un médicament destiné à être utilisé dans le traitement des états associés aux douleurs céphaliques.
 - 13. Utilisation d'un composé selon l'une quelconque des revendications 1 à 10 pour préparer un médicament destiné à être utilisé dans le traitement de la migraine, de la céphalée vasculaire de Horton, de l'hémicrânie paroxysmique chronique ou des céphalées associées à des troubles vasculaires, et pour soulager les symptômes qui leur sont associés.
- 14. Composition pharmaceutique qui comprend au moins un composé de formule (1) selon la revendication 1 ou l'un de ses sels ou produits de solvatation pharmaceutiquement acceptables, avec un ou plusieurs véhicules ou excipients harmaceutiquement acceptables.
- Composition pharmaceutique selon la revendication 14, adaptée à une administration par voie orale, parentérale ou intranasale.
- 16. Composition pharmaceutique selon la revendication 14 ou 15, qui est formulée sous une forme posologique unitaire comprenant de 0.1 à 100 mg du principe actif.
 - 17. Procédé pour préparer un composé de formule (I) selon la revendication 1 ou l'un de ses sels ou produits de solvatation pharmaceuniquement acceptables, qui consiste : (A) A réduire un composé de formule (II)

 $R_1R_2NSO_2(CH_2)_2$

dans laquelle R₁, R₂, R₃ et R₄ sont tels que définis dans la revendication 1 :

OII

(B) à condenser une amine de formule R₁R₂NH (dans laquelle R₁ et R₂ sont tels que définis dans la revendication 1) avec un composé de formule (V)

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(dans laquelle R_3 et R_4 sont tels que définis dans la revendication 1), ou un agent d'acylation correspondant, ou un de leurs sels ou dérivés protégés :

(C) à cycliser un composé de formule (VI)

(D) à réduire un composé de formule (X)

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$$R_1R_2NSO_2(CH_2)_2$$
 $NR_3N=CHCH_2$
 NR_4
 NR_4

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dans laquelle R_1 , R_2 , R_3 et R_4 sont tels que définis dans la revendication 1 ;

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(dans laquelle R₁, R₂, R₃ et R₄ sont tels que définis dans la revendication 1);

(E) à soumettre un autre composé de formule (I) à une réaction d'inter-conversion ;

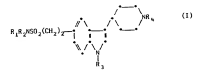
ou

(F) pour préparer un composé de formule (I) dans laquelle P_0 est un groupe alkyle en C_{3-6} , à réduire le composé correspondant dans lequel R_2 représente un groupe alcényle en C_{3-6} ;

- (G) à soumettre un dérivé protégé d'un composé de formule (I) ou d'un de ses sets à une réaction pour éliminer le ou les groupes protecteurs ; et, si cela se révèle nécessaire ou souhaitable, à soumettre le composé obtenu dans l'une quelconque des étapes (A) à (F) à une ou deux réactions supplémentaires consistant :
 - (i) à éliminer des groupes protecteurs,
 - (ii) à convertir un composé de formule (I) ou l'un de ses sels en l'un de ses sels ou produits de solvatation pharmaceutiquement acceptables.

Revendications pour les Etats contractants suivants : GR, ES

1. Procédé pour préparer un composé de formule (I)



dans laquelle

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R₁ représente un atome d'hydrogène ou un groupe alkyle en C₁₋₆ ;

R₂ représente un atome d'hydrogène ou un groupe alkyle en C1-6;

Rs représente un atome d'hydrogène :

R₄ représente un atome d'hydrogène ou un groupe alkyle en C₁₋₃;

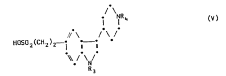
et ses sels et produits de solvatation pharmaceutiquement acceptables, qui consiste :

(A) A réduire un composé de formule (II)

$$R_{1}R_{2}NSO_{2}(CH_{2})_{2}$$

dans laquelle $\mathsf{R}_1,\,\mathsf{R}_2,\,\mathsf{R}_3$ et R_4 sont tels que définis dans la revendication 1 ;

(B) à condenser une amine de formule $R_1\,R_2\,NH$ (dans laquelle R_1 et R_2 sont tels que définis dans la revendication 1) avec un compose de formule (V)



(dans laquelle $\rm H_3$ et $\rm H_4$ sont tels que définis dans la revendication 1), ou un agent d'acylation correspondant, ou un de leurs sels ou dérivés protégés ;

(C) à cycliser un composé de formule (VI)

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$$R_{1}R_{2}NSO_{2}(CH_{2})_{2}-1$$

$$NR_{3}N=CHCH_{2}-1$$

$$NR_{4}$$

$$(VI)$$

dans laquelle R₁, R₂, R₃ et R₄ sont tels que définis dans la revendication 1;

(D) à réduire un composé de formule (X)

$$\mathsf{R}_1\mathsf{R}_2\mathsf{NSO}_2\mathsf{CH}\text{-}\mathsf{CH}\text{-}\bigcup_{\substack{\bullet \\ -\mathbb{N} \\ \mathbb{R}_3}}^{\mathbb{N}^{\mathsf{R}_{\mathsf{R}_{\mathsf{R}}}}} \mathsf{N}^{\mathbb{R}_{\mathsf{R}_{\mathsf{R}}}} \tag{X}$$

(dans laquelle R1, R2, R3 et R4 sont tels que définis dans la revendication 1);

- (E) à soumettre un autre composé de formule (I) à une réaction d'inter-conversion ;
- (F) pour préparer un composé de formule (I) dans laquelle R₂ est un groupe alkyle en C₃-₅, à réduire le composé correspondant dans lequel R₂ représente un groupe alcényle en C₃-₅;
 - (G) à soumettre un dérivé protégé d'un composé de formule (I) ou d'un de ses sels à une réaction pour éliminer le ou les groupes protecteurs ; et, si cela se révèle nécessaire ou souhaitable, à soumettre le composé obtenu dans l'une quelconque des étapes (A) à (F) à une ou deux réactions sunoifémentaires consistant :
 - (i) à éliminer des groupes protecteurs.
 - (ii) à convertir un composé de formule (I) ou l'un de ses sels en l'un de ses sels ou produits de solvatation pharmaceutiquement acceptables.
- Procédé selon la revendication 1, dans lequel R₁ est un atome d'hydrogène ou un groupe alkyle en C₁₋₃.
- 3. Procédé selon la revendication 1 ou 2, dans lequel R₂ est un atome d'hydrogène ou un groupe alkyle
- Procédé selon l'une quelconque des revendications 1 à 3, dans lequel R₂ est un groupe alkyle en C₁-₃.
- 6 5. Procédé selon l'une quelconque des revendications 1 à 4, dans lequel R₁ est un groupe alkyle en C₁₋₂.
 - Procédé selon la revendication 1, dans lequel le produit est le N-méthyl-3-(1-méthyl-4-pipéridinyl)-1Hindole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
 - Procédé selon la revendication 1, dans lequel le produit est le N,N-diméthyl-3-(1-méthyl-4-pipéridinyl)-1H-indole-5-éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.

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- Procédé selon la revendication 1, dans lequel le produit est le N-éthyl-3-(4-pipéridinyl)-1H-indole-5éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
- Procédé selon la revendication 1, dans lequel le produit est le N-méthyl-3-(4-pipéridinyl)-1H-indole-5éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
 - 10. Procédé selon la revendication 1, dans lequel le produit est le 3-(1-méthyl-4-pipéridinyl)-1H-indole-5éthanesulfonamide et ses sels et produits de solvatation pharmaceutiquement acceptables.
- 10 11. Procédé selon l'une quelconque des revendications 1 à 10, dans lequel, dans l'étape (A) ou (D), la réaction est mise en œuvre en présence d'hydrogène et d'un catalyseur à une température de -10 à +50 °C.
- Procédé selon l'une quelconque des revendications 1 à 10, dans lequel, dans l'étape (B), la réaction est mise en œuvre par utilisation d'un agent d'acytation correspondant à la formule (V) à une température de -70 à +150 °C, éventuellement en présence d'une base.
- 13. Procédé selon l'une quelconque des revendications 1 à 10, dans lequel, dans l'étape (C), la réaction est mise en œuvre à une température de 20 à 200 °C.
 - 14. Procédé selon l'une quelconque des revendications 1 à 10, dans lequel, dans l'étape (E), on prépare un composé de formule (I), dans laquelle un ou plusieurs des radicaux R₁, R₂ et R₃ sont des groupes alkyle, par alkylation d'un composé correspondant de formule (I) dans laquelle un ou plusieurs des radicaux R₁, R₂ et R₃ sont des atomes d'hydrogène.
- 15. Procédé pour préparer une composition pharmaceutique, qui consiste à mélanger à un ou plusieurs excipients ou véhicules pharmaceutiquement acceptables un composé de formule (i) selon la revendication 1 ou l'un de ses sels ou produits de solvatation pharmaceutiquement acceptables.

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